Redox stratification of an ancient lake in Gale crater, Mars


INTRODUCTION: The primary goal of NASA's Curiosity rover mission is to explore and quantitatively assess a local region on Mars' surface as a potential habitat for past or present life. A necessary component of that assessment involves an investigation of the surface chemical conditions and paleoclimate of ancient Mars. Gale crater was selected as the landing site for Curiosity; it hosts a ~5-km-tall mountain of layered sedimentary rock. The rocks of Mount Sharp preserve a long-duration record of martian environmental conditions. Geological reconstructions from Curiosity rover data have revealed an ancient, habitable lake environment that was sustained for tens of thousands to tens of millions of years by rivers draining into the crater.

RATIONALE: We seek to constrain the chemical environment within the lake in Gale crater, as well as short- and long-term climate variations in and around Gale crater. We focus on fine-grained sedimentary rocks that carry information about sediment provenance, the environment of deposition, the conversion of sediment to rock during burial (i.e., lithification), and the chemical conditions of later modification (i.e., diagenesis). These were investigated during the first 1300 martian solar days (sols) of rover operations in Gale crater using bulk geochemical and mineralogical analysis techniques, combined with high-resolution color imagery at a variety of scales.

RESULTS: Two mudstone units have been recognized, both deposited in lakes: the Sheepbed member of the Yellowknife Bay formation, an older set of strata defining the base of the stratigraphic section; and the Murray formation, of relatively younger age and positioned higher in the stratigraphic section. The chemical index of alteration (CIA) paleoclimate proxy increases by up to ~10 to 20 CIA units (expressed in %) from the Sheepbed member to the Murray formation. On the basis of mineralogy, geochemistry, textural properties, and stratigraphic relationships, the Murray formation can be subdivided into two sedimentary associations, or facies: the hematite-phyllosilicate (HP) facies and the magnetite-silica (MS) facies. The HP facies is characterized by abundant Fe(III) oxides accompanied by phyllosilicates, as well as indications of Mn oxidation and trace metal concentrations. These properties are consistent with deposition in an oxidizing environment. The MS facies is recognized by a near-complete absence of pure Fe(III) minerals, and high concentrations of silica accompanied by magnetite, consistent with deposition in an anoxic environment. Both facies were affected by a saline overprint after burial and lithification.

CONCLUSION: The observed variations in CIA are consistent with modest short-term fluctuations in the ancient climate between cold, dry conditions and relatively warmer, wetter conditions. These changes occurred during the deposition of lake-bed mudstones in an environment that was conducive to the presence of a long-lived lake in Gale crater. We propose that the distinct properties of the two Murray facies were developed as a result of (i) fractionation of river-borne detritus into coarser, denser materials in shallow water close to shore and finer, lower density material offshore in deeper water as a result of deceleration of river flow as it entered the lake; and (ii) redox stratification of the lake water body, caused by depth-dependent variations in the concentration of atmospheric oxidants and dissolved, groundwater-derived solutes, resulting in oxidizing conditions in shallow water and anoxic in deeper water. The addition of saline minerals during a later phase of brine migration through the section records long-term changes in martian climate at Gale crater, perhaps driven by global atmospheric escape processes. The recognition of redox stratification in the lake in Gale crater adds new detail to our understanding of ancient martian aquatic environments. Previously reported detections of organic carbon compounds, nitrogen, phosphate minerals, and Fe and S minerals in a variety of redox states, combined with the evidence presented here for relatively stable climate conditions and gradients in fluid oxidation state, provide compelling evidence that all of the physical, chemical, and energetic conditions necessary to establish a habitable environment were present on Mars between ~3.8 billion and 3.1 billion years ago. □

The list of author affiliations is available in the full article online.
*Corresponding author. Email: jhurowitz@citytech.cuny.edu

Cite this article as J. A. Hurowitz et al., Science 356, eaah6849 (2017). DOI: 10.1126/science.aah6849
Redox stratification of an ancient lake in Gale crater, Mars

J. A. Hurowitz,1,6 J. P. Grotzinger,2 W. W. Fischer,2 S. M. McLennan,1 R. E. Milliken,3 N. Stein,4 A. R. Vasavada,4 D. F. Blake,5 E. Dehouck,6 J. L. Eigenbrode,7 A. G. Fairén,6,8,9 J. Frydenv10,11,12 R. Geller,11 J. A. Grant,13 S. Gupta,14 K. E. Herkenhoff,15 D. W. Ming,16 E. B. Rampe,17 M. E. Schmidt,18 K. L. Siebach,1,2 K. Stack-Morgan,1 D. Y. Sumner,19 R. C. Wiens10

In 2012, NASA’s Curiosity rover landed on Mars to assess its potential as a habitat for past life and investigate the paleoclimate record preserved by sedimentary rocks inside the ~150-kilometer-diameter Gale impact crater. Geological reconstructions from Curiosity rover data have revealed an ancient, habitable lake environment fed by rivers draining into the crater. We synthesize geochemical and mineralogical data from lake-bed mudstones collected during the first 1300 martian solar days of rover operations in Gale. We present evidence for lake redox stratification, established by depth-dependent variations in atmospheric oxidant and dissolved-solute concentrations. Paleoclimate proxy data indicate that a transition from colder to warmer climate conditions is preserved in the stratigraphy. Finally, a late phase of geochemical modification by saline fluids is recognized.

Fine-grained sedimentary rocks deposited in lakes and oceans provide valuable proxy records of changes in Earth’s dynamic surface environment. These rocks have been used to deduce the nature and extent of variations in global climate (1), the evolution of the composition of Earth’s crust (2), and biologically driven changes in the redox state of Earth’s atmosphere (3). Such rocks have the important property of containing detrital components that carry information about sediment provenance and weathering, and secondary minerals that carry information about the environment of deposition and lithification. In cases where postdepositional geochemical, mineralogical, and textural modification (diagenesis) has partly modified primary signals, it is often possible to decipher those changes to reconstruct past events (4, 5). Given their importance to understanding Earth history, locales containing layered sedimentary rocks linked to depositional processes in rivers, lakes, and seas on Mars have also become a focal point for paleoenvironmental studies. Indeed, a primary motivation behind the selection of Gale crater as the landing site for the Mars Science Laboratory (MSL) Curiosity rover mission was the presence of a ~5-km-tall mountain of layered sedimentary rock, hypothesized to contain a long-duration record of secular changes in martian surface environments (6–9).

As it traversed toward Aeolis Mons (informally known as Mount Sharp), Curiosity has studied mudstones, sandstones, and conglomerates of the Bradbury group and younger mudstones and sandstones of the Murray formation (fm.) at the base of Mount Sharp (10, 11). The lake-bas (lacustrine) mudstones of these two genetically related units form the focus of this study. These rocks are separated by ~60 m of stratigraphic thickness but are related by a common depositional setting (Fig. 1), allowing comparison of their chemical and mineralogical properties. This enables an evaluation of the geochemical and environmental history of the ancient lake system in Gale crater, placing constraints on its habitability and paleoclimate. The Murray fm. can be subdivided into two groups (facies): one that is recognized by abundant ferric iron–bearing minerals (e.g., hematite) accompanied by phyllosilicates and another that is recognized by high concentrations of silica minerals accompanied by magnetite. Mudstones of the Murray fm. show abrupt juxtaposition of these mineral facies, which requires either spatial or temporal variation in lake chemistry. Here we propose a model in which delivery of fine-grained detritus by rivers to a redox-stratified lake provides an explanation for the observed properties of the Murray fm. mudstones. The recognition of a stable redox-stratified water body adds important detail to our understanding of the potential for microbial chemosynthetic metabolism within the ~3.8- to 3.1-billion-year-old Gale crater lake system.

Geological setting

Gale crater is situated in river-dissected terrain immediately south of the hemispherical topographic boundary between the cratered southern highlands and smooth northern lowland plains of Mars, known as the martian dichotomy boundary. Previous work shows that the crater formed ~3.8 billion to 3.6 billion years ago (Ga) (9, 12). The central portion of the crater contains sedimentary deposits that form a large mound (Mount Sharp), possibly cored by a central peak (9, 11–13). After formation, Gale crater was infilled with alluvial fan and lacustrine sediments that were deposited and exhumed before 3.3 to 3.1 Ga (9, 10, 12, 14). Curiosity landed on erosional remnants of a thick sequence (~75 m) of mostly fluvial-deltaic (i.e., river and near-shore) sandstones and conglomerates known as the Bradbury group, which contains a basal unit (Yellowknife Bay fm.) composed in part of lacustrine mudstones [Sheepbed member, Fig. 1, (10, 11, 15, 16)]. To the south of Curiosity’s landing site, the Bradbury group interdigitates with, and is ultimately overlain by, lacustrine mudstones of the Murray fm., which is the lowest exposed unit of the Mount Sharp group. Both the Bradbury group and Murray fm. were buried to substantial depth (hundreds to thousands of meters) and then exhumed by eolian (wind) erosion (10). These rocks have been affected by burial diagenesis, including cementation, fracturing, and vein mineralization (10, 11, 17, 18), in addition to early diagenetic textural modifications, including subaqueous shrinkage cracks, preserved gas bubbles, and nodules (10, 11, 19–21). A younger succession of eolian sandstones (Stimson fm.) overlies the Murray fm. as well as the Bradbury group and is separated from these underlying units by an eolian surface (i.e., an unconformity). Figure S1 shows a map of Curiosity’s traverse and major geomorphic waypoints.

Lacustrine deposits of the Sheepbed member and the younger Murray fm. are predominantly composed of fine-grained rock (<64-μm grain size) displaying massive to laminated (millimeter-to-centimeter-scale) bedding, characteristic of deposition under quiescent conditions (10, 11). Intervals of the Murray fm. show features characteristic of episodic reworking of the lake floor and introduction of coarser sediments by prodeltaic or
fluvial transport, including scour-and-drape deposits, thin centimeter-scale coarser beds, and uncommon thin layers of cross-bedded sandstone (10). The lower Murray fm. shows no evidence for exposure of these sediments while they were wet (e.g., desiccation features), suggesting that this portion of the Gale lake system was perennial. Finally, for the lacustrine strata explored so far, characteristic signs of a glacial climate such as dropstones, diamictites, or ice-wedge fills have not been observed (10, 11).

Through martian solar day (sol) 1300, x-ray diffraction (XRD) data (see methods) were obtained from six mudstone samples by the Chemistry and Mineralogy (CheMin) instrument (22) on board the Curiosity rover (Fig. 1). These samples include John Klein (JK), Cumberland (CL), Confidence Hills (CH), Mojave (MJ), Telegraph Peak (TP), and Buckskin (BK). The distinctive mineralogy of these mudstone samples enables their subdivision into three groupings: (i) the Sheepbed member (which includes JK and CL), (ii) the hematite-phyllosilicate (HP) facies of the Murray fm. (which includes CH and MJ), and (iii) the magnetite-silica (MS) facies of the Murray fm. (which includes TP and BK). On the basis of geochemical, stratigraphic, and sedimentological associations, these groupings can be extended to the full suite of mudstone samples that have been imaged and analyzed by the Curiosity instrument payload, but which have not been analyzed by XRD. The mineralogical and geochemical analysis techniques used in this paper are described in the methods, with data tabulated on table S1 and in data file S1.

Mudstone geochemistry

Gale crater mudstones have been influenced by a combination of (i) variation in source rock composition and degree of chemical weathering (i.e., provenance), (ii) physical sorting during transport and deposition, (iii) water-rock reaction during transport and deposition, (iv) redox and pH conditions within the lake, and (v) diagenetic processes. In Figs. 2 to 4, we describe geochemical relationships, derived from data collected with the Alpha Particle X-ray Spectrometer [APXS, methods, (23)], between and within mudstone suites to understand the influence of these different factors on proxy signals for paleoclimate and redox conditions. The chemical index of alteration [CIA (%)] is plotted against SiO2 [weight % (wt %)] in Fig. 2; CIA is calculated as the ratio \(\frac{Al_2O_3}{Al_2O_3 + Na_2O + CaO + K_2O} \times 100\) in molar proportions, and together, CIA and SiO2 provide information on sediment provenance, paleoweathering conditions, and mineral-addition processes. CIA and SiO2 are positively correlated for the Sheepbed member and Murray HP facies, with the Sheepbed member displaying low CIA and SiO2 and the HP facies displaying higher CIA and SiO2. The MS facies occupies a distinctive position on the diagram, displaced from the HP facies toward intermediate CIA and high SiO2. The CIA values calculated here are minima; they have not been corrected for phases that have the effect of lowering CIA when present (e.g., calcite, halite, gypsum). The effect of these phases on CIA can be accounted for, as
Fig. 2. Paleoweathering, provenance, and mineral addition in Gale crater mudstones. CIA (%) versus SiO₂ (wt %) showing climate-induced variations in mudstone geochemistry. Dashed arrowhead lines are mixing vectors between the MS facies sample named Telegraph Peak (Tel. Pk., gray circles), SiO₂, and CaSO₄. The red arrowhead line shows the vectors generated by a two-stage mixing model where Tel. Pk. and SiO₂ are mixed in 50:50 proportions, followed by the addition of CaSO₄ to the mixed composition. Average martian crust (26) is plotted with a green diamond.

<table>
<thead>
<tr>
<th>Facies</th>
<th>HP facies</th>
<th>MS facies</th>
<th>Tel. Pk. (MS)</th>
<th>Sheepbed</th>
</tr>
</thead>
<tbody>
<tr>
<td>SiO₂ (wt %)</td>
<td>45 ± 4%</td>
<td>43 ± 2%</td>
<td>45 ± 2%</td>
<td>43 ± 2%</td>
</tr>
<tr>
<td>CaSO₄ Addition</td>
<td>50%</td>
<td>50%</td>
<td>50%</td>
<td>50%</td>
</tr>
</tbody>
</table>

CIA reflects the degree to which the sources of sedimentary rocks were chemically weathered before physical erosion and during transport to their depositional site (24, 25). The Sheepbed member mudstones exhibit low CIA values that approximate the CIA of average martian crust, which is 37% (26). Accordingly, McLennan et al. (27) argued that these low CIA values indicate that the sources of these mudstones experienced little to no chemical weathering, consistent with sediment generation and transport under cold and/or arid conditions. In contrast, the HP facies exhibits higher CIA values and SiO₂ concentrations relative to those of the Sheepbed member and average martian crust. A reasonable first-order interpretation is that these elevated CIA values indicate a shift in climate toward conditions that allowed chemical weathering processes to modify the sources of the HP facies and increase their CIA values.

The change in CIA between average Sheepbed member mudstone (45 ± 4%) and HP facies mudstone (45 ± 4%) is 11%, and when the CIA values of the lowest SO₃ mudstone analyses are compared, the difference is 16.1% (supplementary text, fig. S2). These differences might reflect a change from cold to temperate climate conditions in the sediment source regions (1, 24). As stated above, the low CIA values of the Sheepbed member are readily interpreted in terms of a cold and/or arid climate (27), implying that climate conditions were comparatively warmer or wetter during deposition of the Murray fm. However, our understanding of the behavior of CIA in basaltic sedimentary systems is more limited in comparison to sediments generated from the felsic sources that dominate Earth's upper continental crust. Terrestrial sedimentary rocks are typically generated from sources that have a CIA value of ~45 to 50% and evolve to higher values (28). Therefore, we conservatively interpret the differences in CIA shown in Fig. 2 in terms of climate variation from colder and drier to warmer and wetter but cannot infer a specific climate condition in the sediment source regions of the Gale mudstones.

The observed correlation between CIA and SiO₂ is also likely related to variations in the extent of chemical weathering in the sediment source regions. Weathering reactions are expected to liberate aqueous silica to solution (29, 30), thereby increasing the silica concentration of fluids transporting detrital sediment toward Gale crater. Ultimately, this silica would have precipitated in the lacustrine environment during and after deposition of detrital materials. Another possible explanation for the observed CIA is that the igneous sources of HP facies detritus were more evolved (felsic) than those of the Sheepbed member. In this case, the SiO₂ content and CIA of the clastic material in the HP facies would naturally be elevated relative to the less evolved (mafic) clastic material in the Sheepbed member. Indeed, a comparison of the crystalline mineralogy of the Sheepbed member and the Murray fm. indicates that the ratio feldspars/(pyroxenes + olivine) is higher for the Murray fm. than the Sheepbed member (table S1 and (31)). In addition, some fluvial sandstones and conglomerates observed along Curiosity’s traverse exhibit felsic characteristics (32-33). However, felsic igneous rocks with CIA values of ~50% (e.g., rhyolites and granites) typically have SiO₂ concentrations of 70 to 75 wt % (36). The SiO₂ content of the HP facies (51 ± 2 wt %), although elevated compared to the Sheepbed member (43 ± 2 wt %), is not high enough for the differences in CIA between the Sheepbed member and the HP facies to be explained entirely by variations in the proportions of mafic and felsic sources. Accordingly, we interpret the observed correlation between CIA and SiO₂ to result predominantly from chemical weathering processes in the sediment source regions.

Even higher SiO₂ concentrations are observed in the Murray MS facies, consistent with high abundances of crystalline and amorphous silica-rich phases observed in XRD data [see below and (30)]. Low to moderate CIA values are also observed. APXS analyses of samples from the target named Telegraph Peak have among the lowest SiO₂ (52 to 54 wt %) and highest CIA (43%) values of the MS facies. This sample therefore makes a reasonable starting composition from which to model the effects of mineral addition on whole-rock chemistry and better understand the variations in APXS data from the MS facies. The addition of pure SiO₂ results in a horizontal vector in Fig. 2 that does not match the scattered negative relationship between CIA and SiO₂. Fracture- and void-filling CaSO₄ phases are a ubiquitous feature of the sedimentary units in Gale crater that appear to have been emplaced during postlithification diagene (11, 17). A modeled mixture between Telegraph Peak and pure CaSO₄ also does not match the chemical variation displayed by the MS facies (but does provide another mechanism by which positive correlations between CIA and SiO₂ could be generated). Although neither the addition of SiO₂ nor CaSO₄ can explain the position of the MS facies in Fig. 2, on their own, we do find that a two-stage addition model, where SiO₂ is first added to Telegraph Peak followed by CaSO₄ addition, provides a reasonable first-order description of the observed bulk chemical variability of the MS facies. The highest silica analyses in the MS facies (Buckskin, LaMoose) require a 50:50 mixture of Telegraph Peak and SiO₂ followed by the addition of ~2 to 7% CaSO₄ to match their positions. Samples with lower SiO₂ contents and higher CIA values than Buckskin require correspondingly lower amounts of silica and CaSO₄ addition. An implication of this model is that the original CIA values of the MS facies were higher than observed but have been depressed by the presence of Ca sulfate in the APXS field of view (also see supplementary text and fig. S2).

The MS facies also exhibits elevated TiO₂ and P₂O₅ concentrations (figs. S3, A and B); possible causes for these elevated concentrations and potential mineralogical hosts of TiO₂ are discussed in the supplementary text.

Figure 3A plots Zn [parts per million (ppm)] against MnO (wt %), and Fig. 3B plots Ni (ppm) against MnO (wt %). This suite of elements illustrates processes of oxidation and associated trace-metal enrichment, as well as trends developed as a result of dilution processes, during deposition in the lake and continuing during diagenesis. MnO and Zn are positively correlated in the Murray HP facies [regression slope 7500 ± 825, r² = 0.69]. The Murray MS facies samples also exhibit correlated behavior but are excluded from the regression.
Fig. 3. Mn oxidation, metal scavenging, and dilution in the Murray fm. (A) Zn (ppm) versus MnO (wt %) showing a linear regression for the HP facies that results from Mn oxidation and scavenging of aqueous Zn by Mn oxides (the outlier samples Ricardo_Raster_1, Ricardo_DRT_1, Kleinberg, and Schwarzrand are excluded from the regression and labeled with arrowhead lines). (B) Ni (ppm) versus MnO (wt %) demonstrating dilution of MnO and trace metals in the MS facies by the addition of SiO2 and CaSO4, with data trending toward the origin. Green diamond is average Martian crust. The red square is target “Palmwag.”

for reasons discussed below. The Sheepbed member exhibits no obvious trend, with MnO content ranging between ~0.2 and 0.4 wt % and Zn concentrations roughly constant for all samples. In Fig. 3B, neither the Murray HP facies nor the Sheepbed member samples exhibit correlated behavior between Ni and MnO. Both suites of samples are generally enriched in Ni relative to average martian crust, but in this case, the HP facies is indistinguishable from the Sheepbed member. The only exception is a suite of analyses on diagenetic concretions from the base of the HP facies at Pahrump Hills (figs. S1 and S4A), which exhibit high Ni concentrations, ranging from 1934 to 4169 ppm, and MnO concentrations ranging from 0.21 to 0.35 wt %. The MS facies is displaced from the HP facies and the Sheepbed member toward the origin.

For the HP facies, the observed correlation between MnO and Zn can be explained by Mn oxidation and precipitation coupled to scavenging and concentration of Zn from the water column or pore waters during deposition. Mn oxides are highly effective trace metal scavengers that are capable of adsorbing dissolved metal cations to their charged surfaces [e.g., (37)]. This process was invoked to explain the high MnO and Zn concentrations observed in fracture-filling materials between sols 600 and 630 at the Kimberley outcrop (fig. S1, (36)). A second process that may have generated the apparent correlation between MnO and Zn is dilution by chemical and/or mineralogical components that are MnO and Zn free; this would drive samples toward the origin shown in Fig. 3A. In particular, MnO and Zn in the MS facies samples may have been diluted by the addition of silica and Ca sulfate discussed above (Fig. 2). An example of dilution is provided by an analysis of a Ca sulfate–vein target named Palmwag (fig. S4B), which exhibits the lowest MnO and Zn concentrations of any Murray fm. analysis. Analyses of the MS facies sample Telegraph Peak show that it has among the highest MnO and Zn and lowest SiO2 and SO3 concentrations in the MS facies, whereas the remaining MS facies samples, which generally exhibit higher SiO2 and SO3, are correspondingly lower in MnO and Zn. These lines of evidence support the hypothesis that dilution played a role in defining the systematics displayed by the MS facies shown in Fig. 3A.

The process of dilution in the MS facies is also illustrated in Fig. 3B. Although the HP facies and Sheepbed member are both enriched in Ni relative to the average crust of Mars, neither displays correlated behavior between Ni and Mn, and the two suites are indistinguishable from one another in terms of Ni and Mn content. The MS facies, in contrast, is correlated in Mn-Ni space, but in this case, the trend is toward the origin, consistent with a process of dilution by a component that has little to no Mn, Ni, or Zn in it, most likely silica and Ca sulfate. The lack of correlation between MnO and Ni for the HP facies indicates that the process of Mn oxidation and Zn scavenging, hypothesized above, did not have an effect on Ni, despite the fact that both of these metals are expected to be enriched by adsorption on Mn oxides (37). We suggest that the Mn, Zn, and Ni relationships displayed by the HP facies indicate that the primary lake waters from which Mn oxidized and precipitated were relatively Zn rich and Ni poor.

Figure 4 shows wt % MgO versus wt % SO3. These two oxides illustrate the influence of saline fluids on bulk composition. Accordingly, Fig. 4 includes an additional set of samples from the Murray HP facies that are characterized by the presence of sulfate-rich diagenetic concretions, cross-cutting Ca sulfate veins, and/or total SO3 concentrations >9.5 wt % (data file S1). The Sheepbed member exhibits a flat to slightly negative trend in Fig. 4. In contrast, the HP facies samples exhibit a scattered positive linear correlation between MgO and SO3 (slope = 0.33 ± 0.05, r2 = 0.49) when diagenetic and high SO3 samples are included. As a group, the Murray MS facies does not appear to exhibit correlated behavior between MgO and SO3. However, when we exclude analyses of the drill cuttings and tailings piles from the sample Buckskin, which exhibit low MgO (0.5 to 1.9 wt %), the remaining MS facies samples exhibit a weak positive linear correlation (slope = 0.27 ± 0.11, r2 = 0.31), consistent with that exhibited by the HP facies. A suite of analyses of diagenetic concretions in the Stimson fm. (targets Sperrgebiet and Khomas, fig. S4C) provide an additional example of MgO-SO3 correlation. These targets are offset from high total MgO but also exhibit a positive linear correlation between MgO and SO3 with a regression slope (0.31 ± 0.01, r2 = 0.59) that is identical (within error) to that exhibited by the Murray fm. samples. The positive linear correlations observed in Fig. 4 cannot be explained by rock-surface contamination by Gale crater soils, which have an average composition of 8.6 ± 0.9 wt % MgO and 4.6 ± 1.2 wt % SO3 (average ± 1 standard deviation of 10 soil analyses collected between sols 0 and 1184, data file S1), or by atmospheric dust measured at Gale crater, which has identical MgO content to Gale crater soils and SO3 content between 8.0 and 8.3 wt % (39).
higher SiO2 concentrations, our ability to effec-
ponent in these rocks is diluted by increasingly
0.31). Thus, it may be that as the Mg sulfate com-
parison toward high SO3, as shown in Fig. 4. As
would have the effect of depressing the regres-
scales inversely with the SiO2 concentration of
linear correlations between these two oxides
Fig. 4. Magnesium and calcium sulfates in the
Murray fm. MgO (wt %) versus SO3 (wt %), including HP
facies analyses from diagenetic and high SO3
samples (red squares) and diagenetic targets in the
Stimson fm. (Khomass and Sperrgebiet, yellow diamonds).
HP facies and Stimson fm. targets exhibit linear
correlations with slopes of 0.33 ± 0.05 and 0.31 ± 0.01,
respectively. When low-MgO analyses of drill tailings from
the target Buckskin are excluded, the MS facies also
exhibits a positive correlation with slope 0.27 ± 0.11. The
slopes of the regression lines are lower than 0.5, as expected
for Mg sulfate, and consistent with a mixture of Mg- and
Ca-sulfate. Average Gale soil is shown with a green triangle, the green diamond is average Martian
crust, and Sheepbed member is plotted here with small blue circles for clarity.

The positive linear regressions displayed be-
tween MgO and SO3 in Fig. 4 indicate that Mg
sulfate salts play an important role in the HP facies
and possibly in the MS facies as well. In some cases,
these salt enrichments occur as large, centimeter-
scale concretionary or dendritic features ([10, 40],
fig. 5A), but in many cases, they lack a clear tex-
tural expression, suggesting the salts may be a
component of the fine-grained mudstone matrix
or cement. The CheMin instrument has not detect-
ed crystalline Mg sulfate phases, suggesting that
this component is present in an amorphous or
nanocrystalline state. The goodness-of-fit ($r^2$) of
linear correlations between these two oxides
ranges from 0.39 to 0.43 wt % for the Khomas and Sperrgebiet
nodules from the Stimson fm. ($r^2 = 0.99$), from 0.43 to
0.54 wt % for the HP facies samples ($r^2 = 0.47$), and
from 0.54 to 0.73 wt % for the MS facies samples ($r^2 = 0.31$).
Thus, it may be that the Mg sulfate com-
ponent in these rocks is diluted by increasingly higher SiO2 concentrations, our ability to effec-
tively discern Mg sulfate in a bulk chemical analy-
sis decreases. Finally, the slopes of the linear
regressions for the MS facies (0.27 ± 0.11), HP
facies (0.33 ± 0.05), and Stimson fm. (0.31 ± 0.01)
targets are shallower than that of Mg sulfate min-
erals, which are characterized by an MgO:SO3
weight ratio of 0.5. The shallow slopes can be
explained by the incorporation of CaSO4, which
would have the effect of depressing the regression
toward high SO3, as shown in Fig. 4. As
evidence, two samples from the HP facies (San
Francisquito and Palmwag), which intention-
tally targeted cross-cutting Ca sulfate veins, trend
away from the HP samples toward lower MgO
and higher SO3 concentrations.

**Mudstone mineralogy**

In Fig. 5, we describe mineralogical relationships
derived from data collected using the CheMin
instrument to further constrain the nature of
sediment source regions and chemical condi-
tions (e.g., pH, redox state) within the lake. We
tabulated mineral abundances (Fig. 5A, table S1)
in four groups: salts, amorphous, clastic, and prob-
able secondary and redox-sensitive mineral phas-
es. “Salts,” detected at low concentrations (<5 wt %)
in the Sheepbed member and in the MS facies
sample Buckskin, primarily represent detections
of the Ca sulfate minerals anhydrite and bassa-
nite. These fracture-filling materials were inten-
tionally avoided during drill-sample acquisitions,
so their low abundances are expected. “Amorphous”
is an x-ray amorphous component with variable
chemical composition ([18, 31, 41], present at 27
to 54 wt % abundance. “Clastic,” which varies be-
tween ~40 and 50 wt % abundance, includes the
following detrital crystalline minerals of prob-
able igneous origin: feldspars, pyroxenes, olivine,
quartz, cristobalite, tridymite, ilmenite, and apatite.
The “probable secondary and redox-sensitive
mineral phases” group (Fig. 5B, table S1) makes up
the remaining ~10 to 25 wt % of phases
detected by CheMin. This group includes opal-
cristobalite-tridymite (CT), phyllosilicates, the ferric
sulfate mineral jarosite, Fe sulfides (mainly pyr-
rhodite of probable detrital origin), ferric iron
oxides (hematite in the Murray fm., akaganeite
and lesser hematite in the Sheepbed member),
and the mixed-valence iron oxide magnetite.
In the Sheepbed member, this group is dominated
by phyllosilicate (specifically Fe-Mg smectite,
[18, 42]), with lesser magnetite, ferric oxides,
and Fe sulfides, in decreasing order of abundance. The
HP facies samples Confidence Hills and Mojave
contain phyllosilicate, most likely a 2:1-layer phyl-
osilicate such as a collapsed smectite or illite (41),
at lower abundance than in the Sheepbed mem-
er. The Confidence Hills sample also contains
hematite, magnetite, and jarosite, in decreasing
order of abundance; these four phases are pre-
sent in subequal abundances in Mojave. The MS
facies is largely composed of opal-CT and mag-
netite, with minor hematite and jarosite in the
Telegraph Peak sample. No phyllosilicates are ob-
served in the MS facies.

The grouping of mineral and amorphous com-
ponents illustrates processes that influenced the
mineralogical composition of mudstones in Gale
crater (Fig. 5A). A large proportion of all mud-
stones is made up of crystalline minerals of likely
igneous origin that were transported by fluvial
systems as clastic sediment. If we allow for the
possibility that the amorphous component is also
detrital in origin, noting that it has been inter-
preted as a mixture of volcanic glasses and sec-
ondary phases plausibly formed by chemical altera-
tion of that glass ([18, 31, 41, 43], and combine the
amorphous and crystalline group abundances,
then the total ranges from ~70 to 90 wt %. This
large fraction of detrital igneous components
stands in contrast to measurements of terrestrial
Phanerozoic mudrocks, which contain on average
35 to 40 wt % detrital igneous minerals (predom-
nantly quartz and feldspar), with the balance
made up mostly of recycled sedimentary rock
(44, 45). A lack of active plate tectonics on Mars
would contribute to limited or absent sediment
recycling at Gale crater (46). Therefore, we suggest
that the high abundance of detrital primary igneous
minerals and amorphous phases in the mudstones of
Gale crater reflect that they are first-cycle sediments
that directly sample their igneous provenance(s).

Variations displayed by the “probable second-
ary and redox-sensitive mineral phases” group
(Fig. 5B) indicate considerable differences in
fluid chemical conditions during deposition of
the mudstone sample suites. Sheepbed mineralogy
is thought to reflect smectite and magnetite crys-
talization during sediment deposition and burial
(i.e., during authigenesis), resulting from water-
rock reactions with detrital clastic phases under
moderate to alkaline pH, and anoxic to poorly ox-
idizing conditions ([18, 27, 47]. The presence of
detrital pyrrhotite (plus minor pyrite) supports
these conclusions regarding oxidation state, as
do experiments indicating that authigenic mag-
netite can readily form from the spontaneous dis-
proportionation of ferrous hydroxide precipitated
under anoxic, alkaline conditions (48, 49).
The generation of akaganeite might reflect redox dis-
equilibrium in the primary fluids from which
magnetite and smectite precipitated, a common
feature in terrestrial basaltic aquifers (50), or might
record later diagenetic modification by more ox-
idizing fluids.

Further constraints on Gale lake chemistry are
provided by the Murray fm., which shows intern-
al stratigraphic variations in the secondary and
redox-sensitive mineral assemblage. The HP facies
contains smectitic or illitic clay, and the elevated
CIA values observed for the HP facies (Fig. 2) indi-
cate that chemical-weathering reactions were
Hurowitz et al., Science 356, eaah6849 (2017)  2 June 2017

more active in the source regions of the HP facies. Accordingly, some or all of the phyllosilicate in the HP facies may be detrital, reflecting the mineralogy of the sediment source regions (e.g., [31]). Phyllosilicate in the HP facies could also reflect precipitation from neutral to alkaline lake and/or pore waters during authigenesis, similar to the Sheepbed member. The Murray HP facies is characterized by a more oxidized mineral assemblage than the Sheepbed member, with ferric iron-bearing secondary minerals present in greater abundance than mixed-valence iron-bearing species. If we assume that the magnetite in the HP facies is authigenic, similar to what was inferred for the Sheepbed member, then redox disequilibrium or evolving redox conditions during mineral authigenesis are implied. The presence of jarosite in the HP facies requires the activity of acidic fluids, between pH ≈ 2 and 4, which can be readily achieved by the oxidation of sulfide minerals (52). Only ~1.7 wt % detrital pyrrhotite, which is only ~0.5 wt % more than the quantity of detrital pyrrhotite observed in the Sheepbed member, would be required to generate the maximum observed jarosite in the HP facies (3.1 wt % in the Mojave_2 sample). Although CheMin analysis of the HP facies does not indicate the presence of sulfide minerals, the rapid rate of sulfide reaction under oxidizing conditions (33) could readily explain their absence. We suggest that the low-pH conditions implied by small amounts of jarosite do not reflect broader, basin-scale pH conditions but rather were localized to the immediate vicinity of oxidizing fluids in contact with sulfide grains, a phenomenon observed in numerous terrestrial examples (e.g., [54]).

In the MS facies, the total abundance of secondary and redox-sensitive phases varies by more than a factor of 2, and the assemblage is composed almost entirely of opal-CT and magnetite. Possible explanations for the appearance of opaline silica and disappearance of phyllosilicates in the MS facies include the following: (i) Detrital tridymite and cristobalite, present in high abundance in the MS facies, are unstable in surface waters and are expected to decompose to amorphous silica via mineral-water reaction (55, 56). The high abundance of these thermodynamically unstable igneous silica minerals may have favored the production of high-silica pore fluids from which opaline silica precipitated. (ii) Another possibility, suggested by Morris et al. (31), is that opaline silica in the MS facies is detrital in origin. Finally, the MS facies contains extremely low abundances of pure ferric iron-bearing minerals; the near absence of such phases is consistent with anoxic to poorly oxidizing water conditions.

Chemical evolution of the Gale lake system

A simple process model of the lake-basin physiography, sediment transport, and chemistry can be used to explain the observations described above for Murray mudstones (Fig. 6). Fluvial-runoff draining the rim of Gale crater transported clastic detritus ([15, 16]) and fresh water ([18, 27]) to the lake. Gale crater is the lowest point for a thousand kilometers in any direction ([10]), which would have made it a locus for groundwater flow derived from the northward-dipping proximal southern highlands. Therefore, in addition to the freshwater and clastic materials delivered by surface fluvial systems, it is likely that groundwater seepage into the lake bed would have provided an additional source of solutes, such as dissolved SiO₂ (aq), Fe³⁺, Mn²⁺, Zn²⁺, and other ions derived from reactions between water and rock along the regional groundwater flow path. These ions represent the raw materials from which some of the nondenitril secondary minerals could have formed.

Sediment transport from the crater rim and central peak resulted in hydrodynamic sorting that created wedges of fluvial, deltaic, and lacustrine sediments that thinned and fined toward the lake, located in the moat between the crater rim and central peak ([10]). As fluvial waters entered the proximal part of the lake, flow expansion and deceleration deposited thickly laminated sediment and, further offshore, thinly laminated facies representing the tail ends of dispersed plumes (fig. S4, D to F). The decelerating sediment plume would also fractionate clastic detritus so that distal parts would be enriched in lower density materials relative to their near-shore equivalents. Measurements of lamination thicknesses in the Murray fm. show that Si concentration increases and Fe concentration generally decreases with decreasing lamination thickness (fig. S5, A and B). Accordingly, the thicker laminations with more mafic mineralogy (table S1) and silica-poor composition of the Murray HP facies is consistent with its deposition in the proximal, shallower portion of the lake. Conversely, the thinner laminations, siliceous mineralogy (table S1), and more silica-rich composition of the Murray MS facies is consistent with its deposition in a more distal, deeper portion of the lake.

The correlation between the physical and mineralogic attributes of the two Murray lacustrine facies, and their reconstructed paleogeography and bathymetry, allows further development of the lake model. In particular, we can explain why the redox-sensitive mineralogy of the shallow-water HP facies is distinctively different from the deeper-water MS facies. We propose that redox stratification of the lake water body provides a reasonable hypothesis for the differences between these two
Fig. 6. A hypothesized redox-stratified lake in Gale crater. Model of physical transport and geochemical processes occurring during deposition of the Murray fm. Fresh water and clastic materials are delivered by overland flow from fluvial systems; additional dissolved solutes enter the lake by groundwater seepage. Deceleration of the incoming river flow produces characteristic changes in bedforms and sedimentary textures. These are described at the bottom of the diagram with examples from Curiosity’s traverse where such bedforms have been observed (fig. S4, D to F). Flow deceleration also results in sedimentation fractionation; coarser, denser clastic materials are deposited closer to shore, whereas finer, less dense clastics travel further into the lake (also see fig. S5, A and B). Redox stratification results from differences in the mass balance of atmospheric oxidants (UV photons, O2) and oxidizable cations (Fe2+, Mn2+), causing redox-sensitive mineral assemblages to vary as a function of lake water depth.

mineralogic facies and describes an abiotic mechanism of redox stratification for this martian lake. The model depends on the depth of penetration of ultraviolet (UV) light and low levels of photochemically generated atmospheric O2 into the water column to establish a depth-dependent boundary between oxidized and anoxic zones (Fig. 6). Estimation of the depth of the redox transition between oxidized and anoxic waters has a non-unique solution that depends on the concentration of dissolved Fe2+ (and Mn2+), the UV-photon flux reaching the lake surface, the effect of water-column turbidity on UV-light attenuation, the partial pressure of atmospheric O2, the kinetics of gas exchange at the lake-atmosphere interface, and mixing with depth. Measurements of deltaic clinoform thicknesses indicate that the water depths of the most proximal facies were on the order of 1 to 4 m (10), but the maximum depths of the lake are unconstrained by Curiosity data.

In the schematic model displayed in Fig. 6, dissolved, oxidizable cations (Fe2+ and Mn2+) are delivered to the lake primarily by groundwater seepage through lake-floor sediments, whereas oxidants penetrate downward from the lake’s upper surface. In the shallow portion of the lake, oxidant concentration exceeds that of dissolved Fe2+ and Mn2+, so these cations are largely oxidized and precipitate as oxidized minerals, e.g., hematite and Mn oxide. The generation of Mn oxide in the shallow, oxidized portion of the lake results in the scavenging and enrichment of Zn2+ described previously. The relatively low average FeO/MnO ratio of the HP facies (56 ± 20) compared to the MS facies (73 ± 19) supports the hypothesis that Mn bypassed the anoxic portion of the lake but was retained in the oxidized part. The presence of low pH waters, conducive to the small amount of jarosite precipitation that is observed in the HP facies, can be accommodated within the framework of shallow-water oxidative processes. Detrital sulfide minerals would have been oxidized after near-shore deposition, producing locally acidic pore-water fluids. Meanwhile, the lake water body was likely characterized by neutral to alkaline pH values conducive to preservation and/or precipitation of the phyllosilicates observed in the HP facies. Below the depth of penetration of atmospheric oxidants, in the deeper portion of the lake, the waters remain largely anoxic, and under these conditions, the precipitation of authigenic magnetite is favored over more oxidized secondary iron minerals. Ferric iron minerals that settled out into the deeper, anoxic part of the lake could also have undergone reduction by dissolved inorganic ferrous-iron complexes [e.g., (57)] or by H2 gas, which is a by-product of authigenic magnetite precipitation (49), with the net result being a paucity of pure ferric minerals in this part of the lake.

Neoproterozoic iron formations, which make a brief appearance at ~710 million years ago in the terrestrial rock record (58), may provide a useful process analog to the lacustrine mudstones of Gale crater. Unlike their ancient Archean and Paleoproterozoic counterparts, it appears that these iron formations have not interacted strongly with organic matter (59). The occurrence of these Fe-rich deposits at a time in Earth’s history that postdates the rise of atmospheric oxygen [e.g., (60)] implies that Neoproterozoic seawater was, at times, capable of transporting and concentrating ferrous iron in solution and that the process responsible for oxidizing iron in seawater may have been a simple redox titration involving atmospheric oxygen ([60, 61], similar in some respects to what we propose for the shallow portions of the lake represented by the Murray HP facies).

A final component to consider in the context of the stratified lake model is the enrichment of the Murray fm. in Mg and Ca sulfate. Sulfate salts, particularly Mg sulfates, are extremely soluble and precipitate from highly concentrated, saline fluids [e.g., (62)]. Such fluids could readily form via evaporative concentration of lake waters, a scenario that bears some similarity to terrestrial salic-saline lake analogs (63–65). However, in the HP facies, where saline solutions seem to have most affected bulk sediment composition, physical sedimentary structures diagnostic of subaerial exposure and evaporation—such as desiccation cracks, prism cracks, curl-up laminae, fenestrae, intrasets and rip-up chips, and centimeter-scale wave oscillation ripples (66–70)—are absent (10). Most notably, primary bedded evaporites and early diagenetic evaporite minerals with displace growth geometry are absent. In the Murray HP facies, the highest concentrations of MgO and SO3 are most closely associated with burial diagenetic features, such as nodules and dendrites (Fig. 3 and fig. S4A). These observations indicate that instead of a primary evaporitic origin, the saline component was added to the Murray fm. during or after burial and lithification. Diagenetic and high-SO3 targets in the HP facies also display correlated behavior in MnO-Zn space (fig. S6). This suggests that the diagenetic process that added MgO and SO3 to the Murray fm. may also have been oxidizing in nature. A potential driver of saline diagenesis can be found higher up in the stratigraphy of Mount Sharp, beneath the unconformity that defines the boundary between the upper and lower part of the Mount Sharp stratigraphy. There, hundreds of meters of stratified, sulfate-bearing rocks are present, having spectral signatures consistent with the presence of the Mg sulfate mineral kieserite (8, 9). It is possible that these sediments, located hundreds of meters above the base of the Murray fm., were deposited during an evaporitic phase in the evolution of the lake in Gale crater. As the fluids in this lake were driven to an increasingly saline state, they would also have become much denser and could have seeped back through the underlying stratigraphy.
The influence of rapid sedimentation on redox

Contrasting the sedimentologic character of the Sheepbed and Murray mudstones allows the influence of sedimentation rate on redox to be assessed. In particular, the Murray mudstones exhibit well developed lamination (fig. S4E), whereas the Sheepbed mudstones are massive, with only a few thin, continuous lenses of possible siltsone present (J1). In terrestrial mudstones, massive bedding often reflects rapid sedimentation, which can take place when deposition occurs in close proximity to the sediment source (J2). Sheepbed mudstone is directly in contact with fluvial sandstones of the Gillespie member (J2), consistent with the possibility of a proximal sediment source. The Sheepbed member is also characterized by numerous early concretions, mineral-filled subaqueous shrinkage cracks, and possible fenestrae, all formed during early diagenesis (J1, 20, 21); the generation of these features might also have acted to disrupt primary bedding fabrics in the Sheepbed member before lithification. The Sheepbed mudstones contain sulfide minerals of probable detrital origin and magnetite of probable authigenic origin, consistent with anoxia during deposition (J8, 47–49). On the basis of these textural and mineralogical observations, we infer that the Sheepbed mudstones were isolated from the influence of atmospheric oxidants as a result of rapid deposition in a near-shore environment, in which sedimentation rates exceeded oxidation rates. As a result, detrital redox-sensitive grains were preserved, and anoxic conditions were promoted in the sediment column during Sheepbed deposition. Given the cold climate conditions implied by low-CIA values in the Sheepbed, it is plausible that sluggish redox kinetics in the cold water that characterized the lake at this time also contributed to the preservation and precipitation of reduced iron- and sulfur-bearing mineral phases (J25). This mechanism of preserving and promoting anoxic conditions during deposition of the shallow-water Sheepbed member is distinct from that of the deeper water Murray M8 facies, where redox stratification of the water column is implicated.

Hesperian climate and habitability

On the basis of the model we have developed (Fig. 6), we can infer a paleoenvironmental evolution from the mudstone stratigraphy in Gale crater: (i) At the base of the section, in the Sheepbed member, mudstones were deposited under cold climate conditions. Near-shore deposition, proximal to fluvial networks supplying water and clastic detritus to the lake, resulted in rapid fallout of fine-grained sediment from suspension and promoted the development of nonlaminated (massive) mudstone texture. The rapid deposition of Sheepbed sediment also acted to isolate it from atmospheric oxidants. Neutral to alkaline pH and anoxic conditions promoted the formation of authigenic phyllosilicates and magnetite. (ii) The stratigraphically higher Murray fm. was deposited under comparatively warmer and wetter climate conditions. Prograding wedges of sediment, fed by fluvial runoff, thinned with increasing transport distance, resulting in the development of finer lamination toward the center of the lake basin and segregation of detrital minerals on the basis of density and grain size. The mineralogy and geochemistry of the Murray fm. are consistent with a neutral-alkaline–P3D, redox-stratified lake, in which oxidized Fe minerals and phyllosilicates were formed in the shallow portion of the lake, and mixed-valence Fe minerals and silica were formed in deeper portions. (iii) Diagenetic mineralization, possibly driven by downward migrating brines developed during a later, evaporative stage of the lake's evolution, imparted a sulfur-rich character to the Murray fm. and likely played a role in emplacing Ca sulfate–rich veins observed throughout the section.

This record from the mudstones of Gale crater challenges and confirms our current understanding of martian climate evolution. The ~100 m of stratigraphy described here indicate a climate with sufficient warmth and humidity to sustain long-lived lakes in Gale crater (J10). How Mars’ climate system could have achieved the conditions we interpret at Gale crater in the early Hesperian is a matter of current debate (J76). In addition, comparison of the Sheepbed member to the stratigraphically higher Murray fm. indicates an evolution from a colder to a warmer climate state, with only a small component of the observed stratigraphy expressing geochemical properties consistent with a cold climate. Perhaps the climate variations apparent in the record of primary lacustrine environments in Gale crater were influenced by short-term variations, driven by impacts, volcanism, or the obliquity cycle (J77, 78). The recognition of a saline diagenetic overprint on the Murray fm. is broadly consistent with the long-held idea that Mars’ climate was driven from an early, wetter state to a later, more arid condition by atmospheric-loss processes (J77, 79). Finally, organic carbon compounds (J80, 81), nitrogen (J82), phosphate minerals, and Fe and S minerals in a variety of redox states (table S1) are present in the sedimentary rocks of Gale crater. Our analysis of these rocks indicates that gradients in lake-water oxidation state were present in the primary lacustrine environment. Taken together, these results provide compelling evidence that the physical, chemical, and energetic conditions necessary to establish a habitable environment were present on Mars between 3.8 and 3.1 Ga.

Methods

We used element-oxide abundance data obtained by the APXS on board the Curiosity rover for samples analyzed between sols 129 and 487 for the Sheepbed member and sols 696 to 1091 and 1157 to 1279 for the Murray fm. All data used in this paper are available on the Planetary Data System (PDS) Geoscience Node (J83) and are reproduced in data file S1 with their reported analytical uncertainties. A complete description of the instrument, and the methods used for calibration and quantification of APXS data, can be found in Gelert et al. (J23), Campbell et al. (J84), and the PDS (J85). A brief description, modified after information publicly available on the PDS (J85), is provided here: The APXS is a contact instrument that uses curium-244 sources to induce particle-induced x-ray emission (PIXE) and x-ray fluorescence (XRF). The x-ray spectra collected from soil and rock targets were used to determine the abundance of major, minor, and trace elements from sodium to bromine. Low-atomic mass (Z) element x-rays stem from the topmost 5 μm of the sample; higher Z elements like Fe are detected from the upper ~50 μm. The APXS is mounted on a turret at the end of the Curiosity rover’s arm and is deployed on selected targets along the rover traverse to determine their elemental composition. The sampled area is about 1.7 cm in diameter when the instrument is in contact with the sample, and APXS spectra represent the average composition over the sampled area.

Mineral abundances measured by the CheMin XRD instrument for the Sheepbed member samples John Klein and Cunningham, and Murray fm. samples Confidence Hills, Mojave, Telegraph Peak, and Buckskin, are derived from the following data sources: the PDS Geosciences Node (J85), Morris et al. (J31), and Rampe et al. (J41). The measurements are reproduced in table S1 with their reported analytical uncertainties. A complete description of the CheMin instrument and the methods used for calibration and quantification of the CheMin data can be found in Blake et al. (J22) and the PDS (J85). A brief description, modified after information publicly available on the PDS (J85) is provided here: The CheMin instrument is located within the body of the Curiosity rover and consists of a funnel for receiving powdered samples, 27 reusable sample cells, an x-ray source, and a cooled charge-coupled device detector. Data from the CheMin instrument are used to determine and quantify the mineralogy of powdered samples delivered by the rover’s arm-mounted sample-acquisition system. Mineral identification uses various diffraction data libraries (e.g., the International Centre for Diffraction Data, Powder Diffraction File database), and pattern processing and phase quantification use a combination of several available software packages (e.g., Jade, TOPAS, FULLPAT). These approaches produce accurate identification and detection for virtually all crystalline phases at abundances greater than ~1 wt %, whereas x-ray amorphous materials have higher detection limits, with poorer accuracy in quantification.

REFERENCES AND NOTES


4. C. Fedo, H. W. Nesbitt, G. Young, Unraveling the effects of potassium metasomatism in sedimentary rocks and paleosols, with implications for paleoweathering conditions and
83. R. Gellert, Mars Science Laboratory Alpha Particle X-Ray Spectrometer RDR data V1.0, MSL-M-APXS-4/5-RDR-V1.0 (NASA Planetary Data System, 2013).
85. D. Vaniman, Mars Science Laboratory Chemistry and Mineralogy RDR data V1.0, MSL-M-CHEMIN-5-RDR-V1.0 (NASA Planetary Data System, 2012).

ACKNOWLEDGMENTS

We are indebted to the MSL project’s engineering and management teams for their efforts in making the mission effective and enhancing science operations. We are also grateful to those MSL team members who participated in tactical and strategic operations, without whom the data presented here could not have been collected. J.A.H. acknowledges support from a subcontract from the NASA Jet Propulsion Laboratory, California Institute of Technology. J.P.G. acknowledges the support of the NASA Astrobiology Institute. A.G.F. was supported by the Project “icyMARS,” European Research Council Starting Grant no. 307496. J.F. acknowledges the support from the Villum Foundation. R.G. and M.E.S. acknowledge support from the Canadian Space Agency, which is responsible for funding the APXS instrument and its operations. Data presented in this paper are archived in the Planetary Data System at http://pds-geosciences.wustl.edu/missions/msl/index.htm. Three anonymous reviewers are acknowledged for their contributions.

SUPPLEMENTARY MATERIALS

www.sciencemag.org/content/356/6341/eaah6849/suppl/DC1
Supplementary Text
Figs. S1 to S6
Table S1
Data File S1
References (87–99)
31 July 2016, accepted 19 April 2017
10.1126/science.aah6849
Redox stratification of an ancient lake in Gale crater, Mars
Science 356 (6341), . [doi: 10.1126/science.aah6849]

Editor's Summary

The depths of an ancient lake on Mars
Gale crater on Mars was once a lake fed by rivers and groundwater. Hurowitz et al. analyzed 3.5 years of data from the Curiosity rover’s exploration of Gale crater to determine the chemical conditions in the ancient lake. Close to the surface, there were plenty of oxidizing agents and rocks formed from large, dense grains, whereas the deeper layers had more reducing agents and were formed from finer material. This redox stratification led to very different environments in different layers, which provides evidence for Martian climate change. The results will aid our understanding of where and when Mars was once habitable.

Science, this issue p. eaah6849